## => d his

(FILE 'HOME' ENTERED AT 11:25:15 ON 26 FEB 2002) SET COST OFF

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Jan Delaval
     FILE 'HCAPLUS' ENTERED AT 11:25:28 ON 26 FEB 2002
                                                               Reference Librarian
                E OSTEOSCREEN/PA, CS
                                                           Biotechnology & Chemical Library
             13 S E3-E12
L1
                                                             CM1 1E07 - 703-308-4498
                E OESTEOSCREEN/PA, CS
                                                               jan.delaval@uspto.gov
                E MUNDY G/AU
L2
            256 S E3, E6, E8-E10
                E GARRETT R/AU
                E GARRETT R/AU
L3
             55 S E3
                E GARRETT ROSS/AU
L4
              7 S E3, E4
                E ROSSINI G/AU
L5
             80 S E3-E16
                E GARRETT I/AU
L6
             53 S E3-E7
L7
            422 S L1-L6
rs
              4 S L7 AND ?PROTEASOM?
     FILE 'REGISTRY' ENTERED AT 11:29:20 ON 26 FEB 2002
L9
              1 S 140879-24-9
     FILE 'HCAPLUS' ENTERED AT 11:30:15 ON 26 FEB 2002
L10
           2944 S L9
L11
           4723 S PROTEASOME OR PROSOME OR (26S OR 26 S) (L) PROTEASE OR IMMUNOPR
             21 S TRICORN () (PROTEASE OR PROTEINASE)
L12
              4 S L7 AND L10-L12
L13
              4 S L8, L13
L14
          11143 S NF (L) KAPPA (L) B
L15
           8204 S NUCLEAR (L) FACTOR (L) KAPPA (L) B
L16
              4 S L7 AND L15, L16
L17
              5 S L14, L17
L18
             21 S EPOXOMICIN# OR EPOXOMYCIN#
L19
             41 S PS341 OR PS 341
L20
L21
             46 S NLVS
                E ALDEHYDE/CT
L22
            166 S E15(L) (PEPTIDE OR PEPTIDYL)
           1055 S E15+NT(L) (PEPTIDE OR PEPTIDYL)
L23
     FILE 'REGISTRY' ENTERED AT 11:40:11 ON 26 FEB 2002
L24
              1 S 6493-05-6
              1 S 133343-34-7
L25
              7 S LACTACYSTIN
L26
              3 S L26 AND C15H24N2O7S
L27
             19 S C15H24N2O7S/MF
L28
L29
              6 S L28 AND NC4/ES AND 1/NR
L30
              3 S L29 NOT (T/ELS OR GLYCINE)
              1 S 134381-21-8
L31
L32
              9 S C28H50N4O7/MF
L33
              3 S L32 AND OC2/ES AND 1/NR
              2 S L33 NOT T/ELS
L34
L35
              1 S 179324-69-7
              1 S C19H25BN4O4/MF AND NC2NC2/ES AND 46.150.18/RID AND 2/NR
L36
L37
              1 S 158442-41-2
L38
             41 S C32H50N4O8/MF
L39
             14 S L38 AND 4/SQL
              3 S L39 AND 46.150.18/RID AND 1/NR
L40
              1 S 193482-49-4
L41
              3 S C28H43IN4O8S/MF
L42
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2 S L42 NOT 125I
             10 S L24, L25, L30, L31, L34, L35, L36, L37, L41, L43
L44
                SEL RN
L45
             16 S E1-E10/CRN
              3 S L45 NOT MXS/CI
L46
L47
             13 S L44, L46
L48
             13 S L45 NOT L47
     FILE 'HCAPLUS' ENTERED AT 11:49:48 ON 26 FEB 2002
           1978 S L47
L49
L50
              5 S L48
L51
            548 S LACTACYSTIN#
L52
           2396 S PTX
L53
           4729 S L19-L22, L49-L52
     FILE 'REGISTRY' ENTERED AT 11:50:44 ON 26 FEB 2002
L54
                STR
L55
              3 S L54
L56
                STR L54
              1 S L56
L57
                SAV L44 GITOMER1/A TEMP
                SAV L45 GITOMER2/A TEMP
                DEL GITOMER1/A
                DEL GITOMER2/A
                SAV TEMP L47 GITOMER1/A
                SAV TEMP L48 GITOMER2/A
                SAV L56 GITOMER3/Q
     FILE 'HCAPLUS' ENTERED AT 11:57:11 ON 26 FEB 2002
L58
              3 S L7 AND L53
L59
              5 S L18, L58
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=> fil hcaplus

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FILE COVERS 1907 - 26 Feb 2002 VOL 136 ISS 9 FILE LAST UPDATED: 25 Feb 2002 (20020225/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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and/or SDIs in the H/Z/CA/CAplus files incorporating CAS Registry Numbers with the P indicator executed between 12/27/01 and 1/23/02 may be incomplete. See the NEWS message on this topic for more information.

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=> d 159 all hitstr tot
    ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2002 ACS
    2001:300537 HCAPLUS
ΑN
DN
    134:331618
    Inhibitors of proteasomal activity for stimulating bone and hair
ΤI
    arowth
    Mundy, Gregory R.; Garrett, Ross I.; Rossini,
ΙN
PA
    Osteoscreen, Inc., USA
    PCT Int. Appl., 57 pp.
SO
    CODEN: PIXXD2
    Patent
DT
LA
    English
IC
    ICM A61K038-06
    ICS A61K038-07; A61K038-13; A61K031-165; A61K031-365; A61K031-4015;
         A61K031-522; A61P019-00; A61P043-00
CC
    63-6 (Pharmaceuticals)
    Section cross-reference(s): 1, 62
FAN.CNT 1
    PATENT NO.
                     KIND DATE
                                         APPLICATION NO. DATE
     ______
                                          -----
                                                          _____
    WO 2001028579 A2
                           20010426
                                          WO 2000-US41360 20001020
PΙ
    WO 2001028579
                     A3
                           20010920
        W: AU, CA, JP
        RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE
PRAI US 1999-421545
                           19991020
                      Α
    US 2000-558973
                           20000425
                     Α
    Compds. that inhibit the activity of NF-.kappa.
AB
    B or inhibit the activity of the proteasome or both
    promote bone formation and hair growth and are thus useful in treating
    osteoporosis, bone fracture or deficiency, primary or secondary
    hyperparathyroidism, periodontal disease or defect, metastatic bone
    disease, osteolytic bone disease, post-plastic surgery, post-prosthetic
    joint surgery, and post-dental implantation; they also stimulate the
    prodn. of hair follicles and are thus useful in stimulating hair growth,
    including hair d., in subject where this is desirable.
    N-carbobenzyol-Ile-Glu-(OtBu)Ala-Leu-CHO (PSI) in 50% propylene glycol,
    10% DMSO, and 40% water was injected daily for 5 days (1mg/kg body
    wt./day) into the s.c. tissue of mice and the tissue was examd. histol. 16
    days later. The no. of hair follicles increased and the downward
    extension of these hair follicles into the dermal tissue was noted, which
    are hallmarks of anagen. There was an obvious increase in size of the
    follicle diam. and the root sheath diam.
    proteasome inhibitor hair bone growth stimulant
ST
IT
    Transcription factors
    RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (I.kappa.B (inhibitor of NF-
        .kappa.B); inhibitors of proteasomal
        activity for stimulating bone and hair growth)
IT
    Periodontium
     Tooth
        (disease; inhibitors of proteasomal activity for stimulating
        bone and hair growth)
IT
    Hair
        (follicle; inhibitors of proteasomal activity for stimulating
```

bone and hair growth)

```
TT
     Bone, disease
        (fracture; inhibitors of proteasomal activity for stimulating
        bone and hair growth)
ΙT
     Hair preparations
        (growth stimulants; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
     Dental materials and appliances
IT
        (implants; inhibitors of proteasomal activity for stimulating
        bone and hair growth)
IT
     Bone formation
        (inhibitors of proteasomal activity for stimulating bone and
        hair growth)
     Bone morphogenetic proteins
TΤ
     Estrogens
     Growth factors, animal
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (inhibitors of proteasomal activity for stimulating bone and
        hair growth)
IT
     Bone, disease
        (metastatic and osteolytic; inhibitors of proteasomal
        activity for stimulating bone and hair growth)
     Growth factors, animal
ΤT
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (osteogenins; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Surgery
        (post-plastic; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
TΤ
     Hyperparathyroidism
        (secondary; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Phosphoproteins
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (statins; inhibitors of proteasomal activity for stimulating
        bone and hair growth)
ΙT
     Joint, anatomical
        (surgery of; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Osteoporosis
        (therapeutic agents; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
TT
     13598-36-2D, Phosphonic acid, alkylidenebis- derivs.
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (bisphosphonate; inhibitors of proteasomal activity for
        stimulating bone and hair growth)
                          404-86-4, Capsaicin 6493-05-6, PTX
IT
     67-99-2, Gliotoxin
                                   25769-03-3, PDTC
     9035-81-8, Trypsin inhibitor
                                                      59865-13-3, Cyclosporin
                              79902-63-9, Simvastatin 110044-82-1
         65240-86-0, PPM 18
     110115-07-6 133343-34-7, Lactacystin
                                           133407-82-6, MG
           133407-86-0, MG 115 134381-21-8, Epoxomicin
     158442-41-2D, PSI, epoxides
                                   179324-22-2, MG 262
     179324-69-7, PS 341
                           336099-20-8
                   336608-38-9, Bay 11-7082
     336099-21-9
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (inhibitors of proteasomal activity for stimulating bone and
        hair growth)
     140879-24-9, Proteasome
ΙT
```

RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors; inhibitors of **proteasomal** activity for stimulating bone and hair growth)

IT 6493-05-6, PTX 133343-34-7, Lactacystin 134381-21-8, Epoxomicin 158442-41-2D, PSI, epoxides 179324-69-7, PS 341

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (inhibitors of **proteasomal** activity for stimulating bone and hair growth)

RN 6493-05-6 HCAPLUS

CN 1H-Purine-2,6-dione, 3,7-dihydro-3,7-dimethyl-1-(5-oxohexyl)- (9CI) (CA INDEX NAME)

RN 133343-34-7 HCAPLUS

CN L-Cysteine, N-acetyl-, (2R,3S,4R)-3-hydroxy-2-[(1S)-1-hydroxy-2-methylpropyl]-4-methyl-5-oxo-2-pyrrolidinecarboxylate (ester) (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 134381-21-8 HCAPLUS

CN L-Threoninamide, N-acetyl-N-methyl-L-isoleucyl-L-isoleucyl-N-[(1S)-3-methyl-1-[[(2R)-2-methyloxiranyl]carbonyl]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 158442-41-2 HCAPLUS

CN L-Alaninamide, N-[(phenylmethoxy)carbonyl]-L-isoleucyl-L-.alpha.-glutamyl-N-[(1S)-1-formyl-3-methylbutyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 179324-69-7 HCAPLUS

CN Boronic acid, [(1R)-3-methyl-1-[[(2S)-1-oxo-3-phenyl-2-[(pyrazinylcarbonyl)amino]propyl]amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

## IT 140879-24-9, Proteasome

RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors; inhibitors of **proteasomal** activity for stimulating bone and hair growth)

RN 140879-24-9 HCAPLUS

CN Proteinase, multicatalytic (9CI) (CA INDEX NAME)

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*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
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- L59 ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2002 ACS
- AN 2001:240712 HCAPLUS
- DN 135:18367
- TI Therapeutic efficacy of a soluble receptor activator of nuclear factor .kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in a model of humoral hypercalcemia of malignancy
- AU Oyajobi, Babatunde O.; Anderson, Dirk M.; Traianedes, Kathy; Williams, Paul J.; Yoneda, Toshiyuki; Mundy, Gregory R.
- CS Division of Endocrinology, Department of Medicine, University of Texas Health Science Center at San Antonio, San Antonio, TX, 78229, USA
- SO Cancer Res. (2001), 61(6), 2572-2578 CODEN: CNREA8; ISSN: 0008-5472
- PB American Association for Cancer Research
- DT Journal
- LA English
- CC 15-5 (Immunochemistry)
- Receptor activator of NF-.kappa.B (RANK) is AΒ a membrane-bound tumor necrosis factor receptor homolog that mediates signals obligatory for osteoclastogenesis as well as osteoclast activation and survival in vivo. The present study was undertaken to evaluate the efficacy of a sol. murine RANK-human Ig fusion protein (muRANK.Fc) as a bone resorption inhibitor in vitro and in vivo. The in vitro studies demonstrated the ability of muRANK.Fc to inhibit human parathyroid hormone-related protein (PTHrP)-induced resorption in fetal rat long bone cultures. Short-term administration of muRANK.Fc to normal growing mice resulted in a complete disappearance of osteoclasts from metaphyses of long bones assocd. with a pronounced increase in calcified trabeculae and bone radiodensity. In a model of humoral hypercalcemia of malignancy in which PTHrP secreted by s.c. xenografts of human lung cancer in nude mice induces extensive osteolysis and severe hypercalcemia, daily administration of muRANK.Fc from time of tumor implantation profoundly inhibited osteoclastic bone resorption and prevented hypercalcemia. MuRANK.Fc had no effect on tumor prodn. of PTHrP, because there was no difference between circulating human PTHrP levels in muRANK.Fc-treated and vehicle-treated tumor-bearing mice. Moreover, even when treatment was initiated after hypercalcemia was established, muRANK.Fc attenuated further increases in blood ionized calcium. These data demonstrate the potent anti-resorptive effects of muRANK.Fc in vivo as well as highlight the potential utility of disrupting RANK signaling as a novel therapeutic approach in humoral hypercalcemia of malignancy and possibly multiple myeloma and skeletal metastases assocd. with osteolysis.
- ST RANK IgG Fc fusion protein bone resorption hypercalcemia malignancy
- IT Immunoglobulins
  - RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
    - (G, Fc, fusion protein contg.; therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc fusion
    - protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model)
- IT Proteins, specific or class
  - RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(RANK, sol., fusion protein contg.; therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc

fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model)

IT Neoplasm

(humoral hypercalcemia of malignancy; therapeutic efficacy of sol.

receptor activator of NF-.kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model) ΙT Osteoclast (inhibition; therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model) IT(resorption, inhibitors; therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model) IT Signal transduction, biological (therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model) IT 7440-70-2, Calcium, biological studies RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (hypercalcemia; therapeutic efficacy of sol. receptor activator of NF-.kappa.B-IgG Fc fusion protein in suppressing bone resorption and hypercalcemia in humoral hypercalcemia of malignancy model) RE.CNT THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD RE (1) Akatsu, T; Bone 1998, V23, P495 HCAPLUS (2) Anderson, D; Nature 1997, V390, P175 HCAPLUS (3) Body, J; Cancer 2000, V12(Suppl), P3054 (4) Bucay, N; Genes Dev 1998, V12, P1260 HCAPLUS (5) Burgess, T; J Cell Biol 1999, V145, P527 HCAPLUS (6) Capparelli, C; Cancer Res 2000, V60, P783 HCAPLUS (7) Cruz, J; Blood 1998, V92(Suppl 1), P275b (8) Dallas, S; Blood 1999, V93, P1697 HCAPLUS (9) Dougall, W; Genes Dev 1999, V13, P2412 HCAPLUS (10) Emery, J; J Biol Chem 1998, V273, P14363 HCAPLUS (11) Fuller, K; J Exp Med 1998, V188, P997 HCAPLUS (12) Guise, T; Endocr Rev 1998, V19, P18 MEDLINE (13) Guise, T; J Clin Endocrinol Metab 1993, V77, P40 HCAPLUS (14) Hofbauer, L; J Bone Miner Res 2000, V15, P2 HCAPLUS (15) Hofbauer, L; J Clin Endocrinol Metab 2000, V85, P2355 HCAPLUS (16) Horwood, N; Endocrinology 1998, V139, P4743 HCAPLUS (17) Hsu, H; Proc Natl Acad Sci USA 1999, V96, P3540 HCAPLUS (18) Hughes, D; J Bone Miner Res 1995, V10, P1478 HCAPLUS (19) Jimi, E; J Immunol 1999, V163, P434 HCAPLUS (20) Kong, Y; Nature 1999, V397, P315 HCAPLUS (21) Lacey, D; Cell 1998, V93, P165 HCAPLUS (22) Li, J; Proc Natl Acad Sci USA 2000, V97, P1566 HCAPLUS (23) Michigami, T; J Bone Miner Res 1997, V12(Suppl 1), PS106 (24) Moseley, J; Proc Natl Acad Sci USA 1987, V84, P5048 HCAPLUS (25) Mundy, C; Am J Med 1997, V103, P134 (26) Mundy, G; Methods Enzymol 1991, V198, P502 HCAPLUS (27) Nagai, M; Biochem Biophys Res Commun 2000, V269, P523 (28) Oyajobi, B; Bone 1998, V23(Suppl), PS192 (29) Oyajobi, B; J Bone Miner Res 2000, V15(Suppl 1), PS176 (30) O'Brien, C; J Biol Chem 1999, V274, P19310 (31) Pearse, R; Blood 2000, V96(Suppl 1), P549a (32) Pecherstorfer, M; J Bone Miner Res 2000, V15, P147 HCAPLUS (33) Simonet, W; Cell 1997, V89, P309 HCAPLUS

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(35) Suda, T; Endocr Rev 1999, V20, P345 HCAPLUS

(34) Strewler, G; J Clin Investig 1987, V80, P1803 HCAPLUS

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(38) Yasuda, H; Proc Natl Acad Sci USA 1998, V95, P3597 HCAPLUS
    ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2002 ACS
L59
AN
     2000:741943 HCAPLUS
DN
     133:291099
ΤI
     Treatment of myeloma bone disease with proteasomal and
     NF-.kappa.B activity inhibitors
     Mundy, Gregory R.
ΙN
PA
     Osteoscreen, Inc., USA
     PCT Int. Appl., 22 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
IC
     ICM A61K038-04
     ICS A61K031-40; A61K031-166; A61P019-08
CC
     1-6 (Pharmacology)
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                                           ______
                            20001019
                                           WO 2000-US9121
                                                            20000407
PΙ
     WO 2000061167
                      A2
                    · A3
     WO 2000061167
                            20010111
         W: AU, CA, JP
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
                            20020109
                                           EP 2000-921764
                                                            20000407
     EP 1169049
                       Α2
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
PRAI US 1999-289229
                       Α
                            19990409
     WO 2000-US9121
                       W
                            20000407
     The present invention involves the identification and use of compns. for
AB
     treating myeloma bone disease. The compns. inhibit proteasomal
     activity and decrease the activity of the transcription factor NF
     -.kappa.B. Assessment of a candidate compd. for its
     ability to inhibit prodn. or activity of proteasomal enzymes or
     NF-.kappa.B provides a useful means to
     identify agents to treat myeloma bone disease.
     bone myeloma therapy proteasome NFkappaB inhibitor;
ST
     proteasome inhibitor bone myeloma therapy; NF kappaB inhibitor
     bone myeloma therapy
ΙT
     Transcription factors
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (NF-.kappa.B (nuclear
        factor .kappa.B); treatment of myeloma bone
        disease with proteasomal and NF-.kappa.
        B activity inhibitors)
IT
     Antitumor agents
        (multiple myeloma; treatment of myeloma bone disease with
        proteasomal and NF-.kappa.B
        activity inhibitors)
                 65240-86-0, Ppm-18 158442-41-2
TΤ
     5108-96-3
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (treatment of myeloma bone disease with proteasomal and
        NF-.kappa.B activity inhibitors)
ΤТ
     140879-24-9, Proteasome
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (treatment of myeloma bone disease with proteasomal and
        NF-.kappa.B activity inhibitors)
     158442-41-2
TT
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (treatment of myeloma bone disease with proteasomal and
        NF-.kappa.B activity inhibitors)
```

RN 158442-41-2 HCAPLUS

CN L-Alaninamide, N-[(phenylmethoxy)carbonyl]-L-isoleucyl-L-.alpha.-glutamyl-N-[(1S)-1-formyl-3-methylbutyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 140879-24-9, Proteasome

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (treatment of myeloma bone disease with **proteasomal** and

NF-.kappa.B activity inhibitors)

RN 140879-24-9 HCAPLUS

CN Proteinase, multicatalytic (9CI) (CA INDEX NAME)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

L59 ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2002 ACS

AN 2000:478627 HCAPLUS

DN 133:247623

TI Patterns of gene expression associated with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A

AU Ji, Xiaohui; Chen, Di; Xu, Chi; Harris, Steve E.; Mundy, Gregory R.; Yoneda, Toshiyuki

CS Division of Endocrinology and Metabolism, Department of Medicine, University of Texas Health Science Center at San Antonio, San Antonio, TX, USA

SO J. Bone Miner. Metab. (2000), 18(3), 132-139 CODEN: JBMME4; ISSN: 0914-8779

PB Springer-Verlag Tokyo

DT Journal

LA English

CC 2-10 (Mammalian Hormones)

The pluripotent mesenchymal stem cells give rise to osteoblasts, AB adipocytes, chondrocytes, and myoblasts. The differentiation of these stem cells into each of the mature functional cells may be controlled by a distinctive master gene(s) and is assocd. with temporal and spatial expression of diverse genes. Identification of genes that are expressed during the differentiation of the mesenchymal cells to osteoblasts is, therefore, important to obtain insights into the mol. mechanisms of osteogenesis. The murine undifferentiated mesenchymal cell 3T3-F442A, when treated with the bone morphogenetic protein 2 (BMP-2), a well-characterized inducer of mesenchymal cell differentiation, exhibited both osteoblastic and adipocytic differentiation. Using the SAGE (serial anal. of gene expression) technique, which has been shown to enable quant. anal. of large nos. of genes in a simple and quick manner, the authors obtained 1600 sequence tags representing 2107 individual nucleotide sequences from control and BMP-2-treated 3T3-F442A cells, resp. By comparing the frequency of tag occurrence, the authors found profiles of up- or downregulated genes assocd. with osteoblast or adipocyte phenotype

such as type I collagen, osteonectin and OSF-2, or C/EBP.beta., aP2, fatty acid synthase, and lipoprotein lipase, resp., in BMP-2-treated 3T3-F442A cells. The authors' data show that BMP-2 induces not only osteoblastic but also adipocytic differentiation in the 3T3-F442A cells. They also show that the 3T3-F442A cells have bipotentials of differentiating toward osteoblasts and adipocytes. The results, therefore, might explain the inverse correlation between trabecular bone vol. and fat vol. in the bone marrow cavity. The results also suggest that the SAGE may be a useful technique that allows a fast and efficient way to generate global and local views of gene expression assocd. With cellular differentiation of the mesenchymal stem cells.

ST BMP2 gene expression osteoblast adipocyte differentiation

IT Bone morphogenetic proteins

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(2; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) Antigens

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(AD1; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) Chaperonins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(ADP ribosylation factor-like protein 2; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Transcription factors

ΙT

ΙT

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(AP-2 (activator protein 2); patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT RNA formation factors

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(C/EBP-.beta. (CCAAT box/enhancer element-binding protein .beta.); patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Transcription factors

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(Cis2; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT G proteins (guanine nucleotide-binding proteins)

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(Gs (adenylate cyclase-stimulating), .alpha.-subunit; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Histones

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(H2A; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) Heat-shock proteins

IT Heat-shock proteins
 RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(HSC73; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (J1; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) IT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (L12; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) TT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (L22; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) IT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (L32; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) ΙT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (L37a; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) ΙT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (L5; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) IT Proteins, specific or class RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (OSF-2 (osteoblast-specific factor-2); patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) IT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (S16; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) ITRibosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (S2, S28; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) Ribosomal proteins IT RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (S24; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) ΙT Ribosomal proteins RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (S29; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A) ITProteins, specific or class RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (TNF-induced protein complex .gamma.; patterns of gene expression

assocd. with BMP-2-induced osteoblast and adipocyte differentiation of

mesenchymal progenitor cell 3T3-F442A)

IT Phosphoproteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(acidic ribosomal protein P2; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Phosphoproteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(acidic ribosomal, P1; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Phosphoproteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(acidic ribosomal, PO; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Phosphoproteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process) (adducins, human erythrocyte, .alpha.-subunit; patterns of gene

expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Adipose tissue

(adipocyte, differentiation; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Cell differentiation

(adipocyte; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Proteins, specific or class

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(calcylin; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Proteins, specific or class

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(calgizzarins; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Chaperonins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(chaperone CCTB; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Osteoblast

(differentiation; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(human ribosomal protein S20; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological

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study); FORM (Formation, nonpreparative); PROC (Process)
   (human ribosomal protein S7; patterns of gene expression assocd. with
   BMP-2-induced osteoblast and adipocytė differentiation of mesenchymal
   progenitor cell 3T3-F442A)
Proteins, specific or class
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (hydrophobic protein MTF; patterns of gene expression assocd. with
   BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal
   progenitor cell 3T3-F442A)
Proteins, specific or class
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (insulin-stimulated eIF-4E binding protein; patterns of gene expression
   assocd. with BMP-2-induced osteoblast and adipocyte differentiation of
   mesenchymal progenitor cell 3T3-F442A)
Proteins, specific or class
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (jesolin; patterns of gene expression assocd. with BMP-2-induced
   osteoblast and adipocyte differentiation of mesenchymal progenitor cell
   3T3-F442A)
Transcription factors
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (junB; patterns of gene expression assocd. with BMP-2-induced
   osteoblast and adipocyte differentiation of mesenchymal progenitor cell
   3T3-F442A)
Proteins, specific or class
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (minopontins; patterns of gene expression assocd. with BMP-2-induced
   osteoblast and adipocyte differentiation of mesenchymal progenitor cell
   3T3-F442A)
Proteins, specific or class
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (mitochondrial ATPase inhibitor; patterns of gene expression assocd.
   with BMP-2-induced osteoblast and adipocyte differentiation of
   mesenchymal progenitor cell 3T3-F442A)
Cell differentiation
   (osteoblast; patterns of gene expression assocd. with BMP-2-induced
   osteoblast and adipocyte differentiation of mesenchymal progenitor cell
   3T3-F442A)
Transcription factors
RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
study); FORM (Formation, nonpreparative); PROC (Process)
   (p68-c-rel; patterns of gene expression assocd. with BMP-2-induced
   osteoblast and adipocyte differentiation of mesenchymal progenitor cell
   3T3-F442A)
Bone formation
   (patterns of gene expression assocd. with BMP-2-induced osteoblast and
   adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)
Gene, animal
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
   (patterns of gene expression assocd. with BMP-2-induced osteoblast and
   adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)
Chloride channel
Fibroblast growth factor receptors
Macrophage migration inhibitory factor
Osteonectin
Ribosomal proteins
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Tubulins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Proteins, specific or class

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(protein for hereditary multiple exostosis; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Proteins, specific or class

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(rat brain protein; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(rat ribosomal protein L23A; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(rat ribosomal protein S19; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Ribosomal proteins

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(rpA2; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Embryo, animal

(stem cell; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Collagens, biological studies

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(type I; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT Anion channel

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(voltage-dependent 3; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

IT G proteins (guanine nucleotide-binding proteins)

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(.beta.-subunit; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

## IT 140879-24-9, Proteasome

RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(Rc7-I; patterns of gene expression assocd. with BMP-2-induced osteoblast and adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)

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IT
     147014-97-9, CDK4 kinase
     RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
     study); FORM (Formation, nonpreparative); PROC (Process)
        (inhibitor; patterns of gene expression assocd. with BMP-2-induced
        osteoblast and adipocyte differentiation of mesenchymal progenitor cell
        3T3-F442A)
                                     9007-43-6, Cytochrome c, biological
ΙT
     9004-02-8, Lipoprotein lipase
               9036-37-7, Aminolevulinic acid dehydrogenase
                                                              9045-77-6, Fatty
                     9059-25-0, Lysyl oxidase
                                                9059-32-9, GTPase
    acid synthase
    Cathepsin L
     RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
     study); FORM (Formation, nonpreparative); PROC (Process)
        (patterns of gene expression assocd. with BMP-2-induced osteoblast and
        adipocyte differentiation of mesenchymal progenitor cell 3T3-F442A)
ΙT
     9001-16-5, Cytochrome c oxidase
     RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
     study); FORM (Formation, nonpreparative); PROC (Process)
        (subunit VIII; patterns of gene expression assocd. with BMP-2-induced
        osteoblast and adipocyte differentiation of mesenchymal progenitor cell
        3T3-F442A)
     37205-63-3, ATP synthase
TT
    RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
     study); FORM (Formation, nonpreparative); PROC (Process)
        (.gamma.-chain precursor and hydrogen-transporting; patterns of gene
        expression assocd. with BMP-2-induced osteoblast and adipocyte
        differentiation of mesenchymal progenitor cell 3T3-F442A)
             THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD
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IT
    140879-24-9, Proteasome
     RL: BPR (Biological process); MFM (Metabolic formation); BIOL (Biological
     study); FORM (Formation, nonpreparative); PROC (Process)
        (Rc7-I; patterns of gene expression assocd. with BMP-2-induced
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osteoblast and adipocyte differentiation of mesenchymal progenitor cell

3T3-F442A) RN 140879-24-9 HCAPLUS CN Proteinase, multicatalytic (9CI) (CA INDEX NAME) \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\* ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2002 ACS ΑN 2000:53374 HCAPLUS DN 132:102860 Inhibitors of proteasomal activity for stimulating bone and hair ΤI IN Mundy, Gregory R.; Garrett, I. Ross; Rossini, PΑ Osteoscreen, USA PCT Int. Appl., 39 pp. SO CODEN: PIXXD2 DTPatent LA English IC ICM A61K031-00 CC 1-12 (Pharmacology) Section cross-reference(s): 63 FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ---------WO 1999-US15533 19990709 WO 2000002548 A2 20000120 PΤ W: AL, AM, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GE, HU, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, SD, SG, SI, SK, TR, TT, US, UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 20000201 AU 1999-63109 19990709 AU 9963109 A1 EP 1096924 A1 20010509 EP 1999-933827 19990709 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI 19980710 PRAI US 1998-113947 A1 WO 1999-US15533 W 19990709 AB Compds. that inhibit the activity of NF-.kappa. B or inhibit the activity of the proteasome or both promote bone formation and hair growth and are thus useful in treating osteoporosis, bone fracture or deficiency, primary or secondary hyperparathyroidism, periodontal disease or defect, metastatic bone disease, osteolytic bone disease, post-plastic surgery, post-prosthetic joint surgery, and post-dental implantation. They also stimulate the prodn. of hair follicles and are thus useful in stimulating hair growth, including hair d., in subject where this is desirable. hair bone growth stimulation NFkappaB inhibitor; proteasome ST inhibitor hair bone growth stimulation Transcription factors IT RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (NF-.kappa.B (nuclear factor .kappa.B); NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Bone formation Drug delivery systems Drug screening (NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) Bone morphogenetic proteins IT Estrogens

Growth factors, animal Hormones, animal, biological studies RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth, and use with other agents) ΙT Antitumor agents (bone, metastasis; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Skull (calvarium, calvarial bone growth assay; NF-.kappa. B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Cartilage (cartilage-derived morphogenetic proteins; NF-.kappa. B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth, and use with other agents) Joint, anatomical IT (degeneration; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) ΙT Disease, animal (dental; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Periodontium (disease; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Hair (follicle; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) Bone, disease IT (fracture, and bone deficiency; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Bone (growth promoters; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth, and use with other agents) IT Hair preparations (growth stimulants; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Dental materials and appliances (implants, post-dental implantation; NF-.kappa. B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) IT Cell differentiation (inducers; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth, and use with other agents) IT Bone, neoplasm (metastasis, inhibitors; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) Proteins, specific or class IT RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (morphogenetic, cartilage-derived; NF-.kappa.

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B inhibitors and inhibitors of proteasomal activity
        for stimulating bone and hair growth, and use with other agents)
IT
     Growth factors, animal
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (osteogenins; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth, and use with other agents)
ΙT
     Bone, disease
        (osteolytic; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth)
IT
     Isoprenoids
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (pathway; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth)
IT
     Peptides, biological studies
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (peptidic aldehydes; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
TΤ
     Aldehydes, biological studies
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (peptidyl; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Surgery
        (plastic, post-plastic surgery; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
     Joint, anatomical
ΤT
     Prosthetic materials and Prosthetics
        (post-prosthetic joint surgery; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Hyperparathyroidism
        (primary; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth)
     Proteins, specific or class
TΤ
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (proteasome; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IT
     Bone
        (resorption, inhibitors; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth, and use with other agents)
IT
     Hyperparathyroidism
        (secondary; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth)
IT
     Osteoporosis
        (therapeutic agents; NF-.kappa.B
        inhibitors and inhibitors of proteasomal activity for
        stimulating bone and hair growth)
IΤ
     Drug delivery systems
        (topical; NF-.kappa.B inhibitors and
        inhibitors of proteasomal activity for stimulating bone and
        hair growth)
```

67-99-2, Gliotoxin 404-86-4, Capsaicin 6493-05-6, 59865-13-3, Cyclosporin A 79902-63-9, Simvastatin Pentoxifylline 106096-93-9, Basic fibroblast growth factor 110044-82-1 **133343-34-7**, Lactacystin 133407-82-6, MG 132 133407-86-0, MG 115 **158442-41-2** 179324-22-2, MG 262 RL: BAC (Biological activity or effector, except adverse); THU . (Therapeutic use); BIOL (Biological study); USES (Uses) (NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) ΙT 140879-24-9, Proteasome RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) 13598-36-2D, Phosphonic acid, bisphosphonates ΙT RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (and statins; NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth, and use with other agents) 6493-05-6, Pentoxifylline 133343-34-7, IT Lactacystin 158442-41-2 RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (NF-.kappa.B inhibitors and inhibitors of proteasomal activity for stimulating bone and hair growth) RN 6493-05-6 HCAPLUS 1H-Purine-2,6-dione, 3,7-dihydro-3,7-dimethyl-1-(5-oxohexyl)- (9CI) (CA CN INDEX NAME)

RN 133343-34-7 HCAPLUS
CN L-Cysteine, N-acetyl-, (2R,3S,4R)-3-hydroxy-2-[(1S)-1-hydroxy-2methylpropyl]-4-methyl-5-oxo-2-pyrrolidinecarboxylate (ester) (9CI) (CA
INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 158442-41-2 HCAPLUS

CN L-Alaninamide, N-[(phenylmethoxy)carbonyl]-L-isoleucyl-L-.alpha.-glutamyl-N-[(1S)-1-formyl-3-methylbutyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 140879-24-9, Proteasome

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(NF-.kappa.B inhibitors and inhibitors of

proteasomal activity for stimulating bone and hair growth)

RN 140879-24-9 HCAPLUS

CN Proteinase, multicatalytic (9CI) (CA INDEX NAME)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

=> fil biosis

FILE 'BIOSIS' ENTERED AT 16:19:56 ON 26 FEB 2002 COPYRIGHT (C) 2002 BIOLOGICAL ABSTRACTS INC.(R)

FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 21 February 2002 (20020221/ED)

=> d all tot

L193 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2002 ACS

```
AN
     1998:745665 HCAPLUS
     130:94381
DN
ΤI
    NF-.kappa.B activation provides the
     potential link between inflammation and hyperplasia in the
     arthritic joint
ΑU
     Miagkov, Alexei V.; Kovalenko, Dmitry V.; Brown, Chadwick E.; Didsbury,
     John R.; Cogswell, John P.; Stimpson, Stephen A.; Baldwin, Albert S.;
     Makarov, Sergei S.
     Thurston Arthritis Research Center, University of North Carolina, Chapel
CS
     Hill, NC, 27599, USA
     Proc. Natl. Acad. Sci. U. S. A. (1998), 95(23), 13859-13864
SO
     CODEN: PNASA6; ISSN: 0027-8424
     National Academy of Sciences
PΒ
DT
     Journal
LA
     English
CC
     15-8 (Immunochemistry)
     Section cross-reference(s): 3
AB
     The transcription factor NF-.kappa.B is a
     pivotal regulator of inflammatory responses. While the activation of
     NF-.kappa.B in the arthritic
     joint has been assocd. with rheumatoid arthritis (RA), its
     significance is poorly understood. Here, the authors examine the role of
     NF-.kappa.B in animal models of RA. The
     authors demonstrate that in vitro, NF-.kappa.B
     controlled expression of numerous inflammatory mols. in synoviocytes and
     protected cells against tumor necrosis factor .alpha. (TNF.alpha.) and Fas
     ligand (FasL) cytotoxicity. Similar to that obsd. in human RA, NF
     -.kappa.B was activated in the synovium of rats with
     streptococcal cell wall (SCW)-induced arthritis. In vivo suppression of
    NF-.kappa.B by either proteasomal
     inhibitors or intraarticular adenoviral gene transfer of super-repressor
     I.kappa.B.alpha. profoundly enhanced apoptosis in the
     synovium of rats with SCW- and pristane-induced arthritis. This indicated
     that the activation of NF-.kappa.B protected
     the cells in the synovium against apoptosis and thus provided the
     potential link between inflammation and hyperplasia. Intraarticular
     administration of NF-kB decoys prevented the recurrence of SCW
     arthritis in treated joints. Unexpectedly, the severity of arthritis also
     was inhibited significantly in the contralateral, untreated joints,
     indicating beneficial systemic effects of local suppression of NF
     -.kappa.B. These results establish a mechanism
     regulating apoptosis in the arthritic joint and
     indicate the feasibility of therapeutic approaches to RA based on the
     specific suppression of NF-.kappa.B.
     transcription factor NFkappaB cytokine inflammation hyperplasia rheumatoid
ST
     arthritis
     Apoptosis
ΙT
     Rheumatoid arthritis
     Synoviocyte
     Transcriptional activation
        (NF-.kappa.B activation in inflamed
        synovium activates inflammatory cytokines but inhibits TNF.alpha.- and
        FasL-mediated apoptosis thereby promoting hyperplasia in animal models
        of rheumatoid arthritis)
ΙT
     NF-.kappa.B
     RL: BAC (Biological activity or effector, except adverse); BIOL
     (Biological study)
        (NF-.kappa.B activation in inflamed
        synovium activates inflammatory cytokines but inhibits TNF.alpha.- and
        FasL-mediated apoptosis thereby promoting hyperplasia in animal models
        of rheumatoid arthritis)
     Fas ligand
IT
```

Interleukin 1.beta.

ΙT

IT

IT

ΙT

Interleukin 6 Tumor necrosis factor .alpha. VCAM-1 (cell adhesion molecule) RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process) (NF-.kappa.B activation in inflamed synovium activates inflammatory cytokines but inhibits TNF.alpha. - and FasL-mediated apoptosis thereby promoting hyperplasia in animal models of rheumatoid arthritis) Synovial membrane (disease, synovitis; NF-.kappa. B activation in inflamed synovium activates inflammatory cytokines but inhibits TNF.alpha. - and FasL-mediated apoptosis thereby promoting hyperplasia in animal models of rheumatoid arthritis) Synovial membrane (hyperplasia; NF-.kappa.B activation in inflamed synovium activates inflammatory cytokines but inhibits TNF.alpha.- and FasL-mediated apoptosis thereby promoting hyperplasia in animal models of rheumatoid arthritis) Hyperplasia (synovial; NF-.kappa.B activation in inflamed synovium activates inflammatory cytokines but inhibits TNF.alpha. - and FasL-mediated apoptosis thereby promoting hyperplasia in animal models of rheumatoid arthritis) Inflammation (synovitis; NF-.kappa.B activation in inflamed synovium activates inflammatory cytokines but inhibits TNF.alpha. - and FasL-mediated apoptosis thereby promoting hyperplasia in animal models of rheumatoid arthritis) THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT (1) Asahara, H; J Rheumatol 1997, V24, P430 MEDLINE (2) Baldwin, A; Annu Rev Immunol 1996, V14, P649 HCAPLUS (3) Beg, A; Science 1996, V274, P782 HCAPLUS (4) Bielinska, A; Science 1990, V267, P891 (5) Brockman, J; Mol Cell Biol 1995, V15, P2809 MEDLINE (6) Brown, K; Science 1995, V267, P1485 HCAPLUS (7) Cheshire, J; Mol Cell Biol 1993, V17, P6746 (8) Firestein, G; Am J Pathol 1996, V149, P2143 MEDLINE (9) Firestein, G; Arthritis Rheum 1996, V39, P1781 MEDLINE (10) Firestein, G; J Clin Invest 1995, V96, P1631 HCAPLUS (11) Fujisawa, K; Arthritis Rheum 1996, V39, P197 HCAPLUS (12) Ghivizzani, S; Proc Natl Acad Sci USA 1998, V95, P4613 HCAPLUS (13) Handel, M; Arthritis Rheum 1995, V38, P1762 MEDLINE (14) Kaltschmidt, C; Biol Chem Hoppe-Seyler 1995, V376, P9 HCAPLUS (15) Liu, Z; Cell 1996, V87, P565 HCAPLUS (16) Makarov, S; Proc Natl Acad Sci USA 1996, V93, P402 HCAPLUS (17) Marok, R; Arthritis Rheum 1996, V39, P583 HCAPLUS (18) Morishita, R; Nat Med 1997, V3, P894 HCAPLUS (19) Mountz, J; Arthritis Rheum 1994, V37, P1415 MEDLINE (20) Nakajima, T; Arthritis Rheum 1995, V38, P485 MEDLINE (21) Nishioka, K; Arthritis Rheum 1998, V41, P1 MEDLINE (22) Nita, I; Arthritis Rheum 1996, V39, P820 MEDLINE (23) Palombella, V; Cell 1994, V78, P773 HCAPLUS (24) Ponton, A; J Biol Chem 1996, V271, P8991 HCAPLUS (25) Roessler, B; J Clin Invest 1993, V92, P1085 HCAPLUS (26) Schwab, J; Mechanisms and Models In Rheumatoid Arthritis 1995, P431 HCAPLUS (27) van Antwerp, D; Science 1996, V274, P787 HCAPLUS (28) Vingsbo, C; Am J Pathol 1996, V149, P1675 HCAPLUS (29) Wang, C; Science 1996, V274, P784 HCAPLUS

(30) Zhou, T; J Immunol 1996, V156, P2661 HCAPLUS

(31) Zvaifler, N; Am J Pathol 1997, V150, P1125 MEDLINE

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L193 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2002 ACS
     1998:263658 HCAPLUS
DN
     129:15209
TΙ
     Activation of NF-.kappa.B is involved in the
     survival of osteoclasts promoted by interleukin-1
ΑU
     Jimi, Eijiro; Nakamura, Ichiro; Ikebe, Tetsuro; Akiyama, Shuichi;
     Takahashi, Naoyuki; Suda, Tatsuo
     Dep. Biochem., School Dentistry, Showa Univ., Tokyo, 142-8555, Japan
CS
     J. Biol. Chem. (1998), 273(15), 8799-8805
SO
     CODEN: JBCHA3; ISSN: 0021-9258
PB
     American Society for Biochemistry and Molecular Biology
DT
     Journal
LA
     English
CC
     15-5 (Immunochemistry)
AB
     The authors previously reported that interleukin-1 (IL-1) promoted the
     survival of murine osteoclast-like cells (OCLs) formed in vitro
     and activated a transcription factor, NF-.kappa.
     B, of OCLs. The present study examd. whether the activation of
     NF-.kappa.B is directly involved in the
     survival of OCLs promoted by IL-1. The expression of IL-1 type I receptor
     mRNA in OCLs was detected by the PCR amplification of reverse-transcribed
     mRNA. An electrophoretic mobility shift assay showed that IL-1
     transiently activated NF-.kappa.B in the
     nuclei of the OCLs, and the maximal activation occurred at 30 min.
     degrdn. of I.kappa.B.alpha. coincided with the
     activation of NF-.kappa.B in the OCLs. The
     immunocytochem. study revealed that p65, a subunit of NF-.
     kappa.B, was translocated from the cytoplasm into almost
     all of the nuclei of the OCLs within 30 min after IL-1 stimulation. The
     purified OCLs spontaneously died via apoptosis, and IL-1 promoted the
     survival of OCLs by preventing their apoptosis. The pretreatment of
    , purified OCLs with proteasome inhibitors suppressed the
     IL-1-induced activation of NF-.kappa.B and
     prevented the survival of OCLs supported by IL-1. When OCLs were
     pretreated with antisense oligodeoxynucleotides to p65 and p50 of
     NF-.kappa.B, the expression of resp. mRNAs by
     OCLs was suppressed, and the IL-1-induced survival of OCLs was
     concomitantly inhibited. Thus, IL-1 promotes the survival of
     osteoclasts through the activation of NF-.kappa
     .B.
ST
     NF kappaB survival osteoclast interleukin 1
ΙT
     Apoptosis
       Osteoclast
        (interleukin-1 promotes osteoclasts survival by preventing
        apoptosis via NF-.kappa.B activation)
IT
     Interleukin 1
     RL: BAC (Biological activity or effector, except adverse); BIOL
     (Biological study)
        (interleukin-1 promotes osteoclasts survival by preventing
        apoptosis via NF-.kappa.B activation)
IT
     NF-.kappa.B
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (interleukin-1 promotes osteoclasts survival by preventing
        apoptosis via NF-.kappa.B activation)
L193 ANSWER 3 OF 8 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. DUPLICATE 1
     1997:300447 BIOSIS
ΑN
     PREV199799599650
DN
     The trental influence on collagen proteolysis in experimental
ΤI
     aseptic infarction of the long bone.
ΑU
     Magomedov, S.; Grigorovskii, V. V.
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Ukr. Res. Inst. Traumatol. Orthop., Ukr. Minist. Health, Kiev Ukraine

CS

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gitomer ~ 09 / 695807
SO.
     Ukrainskii Biokhimicheskii Zhurnal, (1996) Vol. 68, No. 5, pp. 69-76.
     ISSN: 0201-8470.
     Article
DT
LA
     Russian
SL
     Ukrainian; English
     Dynamics of biochemical parameters of the connective tissue and
AB
     morphometric parameters of lesion were studied in rabbits with induced
     embolic aseptic infraction of the femur without and with the
     trental (pentoxyphyllin) treatment. The correlation was
     found between the pairs of indices: proteolytic activity and bone marrow
     necrosis volume: collagenase activity and bone cortex remodelling rate:
     concentration of protein bound with hydroxyprolin fraction and endosteal
     regenerate volume.
CC
     Biochemical Studies - General *10060
     Cardiovascular System - General; Methods *14501
       Bones, Joints, Fasciae, Connective and Adipose Tissue - General;
     Methods *18001
     Pharmacology - General *22002
               *86040
BC
     Leporidae
ΙT
     Major Concepts
        Biochemistry and Molecular Biophysics; Cardiovascular System (Transport
        and Circulation); Pharmacology; Skeletal System (Movement and Support)
ΙT
    Chemicals & Biochemicals
          TRENTAL; PENTOXIFYLLINE; COLLAGENASE
     Miscellaneous Descriptors
ΙT
        ASEPTIC INFARCTION; BONE CORTEX REMODELLING RATE; BONE DISEASE; BONE
        MARROW NECROSIS VOLUME; COLLAGEN PROTEOLYSIS; COLLAGENASE ACTIVITY;
        ENDOSTEAL REGENERATE VOLUME; EXPERIMENTAL; FEMUR; LONG
        BONE; PENTOXIFYLLINE; PENTOXYPHYLLIN;
        PHARMACOLOGY; SKELETAL SYSTEM; TRENTAL INFLUENCE; VASCULAR
        DISEASE; VASODILATOR-DRUG
ORGN Super Taxa
        Leporidae: Lagomorpha, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        rabbit (Leporidae)
ORGN Organism Superterms
        animals; chordates; lagomorphs; mammals; nonhuman mammals; nonhuman
        vertebrates; vertebrates
RN
     6493-05-6 (TRENTAL)
       6493-05-6 (PENTOXIFYLLINE)
     9001-12-1 (COLLAGENASE)
L193 ANSWER 4 OF 8 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
ΑN
     2001:561734 BIOSIS
DN
     PREV200100561734
     Regulation of osteoblast differentiation by proteasome control
ΤI
     of Smadl.
ΑU
     Chen, D. (1); Zhao, M. (1); Qiao, M. (1); Garrett, R. (1); Mi,
     Z. (1); Crews, C.; Mundy, G. (1)
     (1) Medicine, University of Texas Health Science Center at San Antonio,
CS
     San Antonio, TX USA
     Journal of Bone and Mineral Research, (September, 2001) Vol. 16, No.
SO
     Suppl. 1, pp. S145. print.
     Meeting Info.: Twenty-Third Annual Meeting of the American Society for
     Bone and Mineral Research Phoenix, Arizona, USA October 12-16, 2001
     ISSN: 0884-0431.
DT
     Conference
LA
     English
```

- SL English
- CC General Biology Symposia, Transactions and Proceedings of Conferences, Congresses, Review Annuals \*00520
  Cytology and Cytochemistry Animal \*02506
  Bones, Joints, Fasciae, Connective and Adipose Tissue Physiology and

Biochemistry \*18004 Animalia - Unspecified 33000 BC IT Major Concepts Skeletal System (Movement and Support) IT Chemicals & Biochemicals Smad-1 protein: osteoblast differentiation regulator, proteasome control Miscellaneous Descriptors IT Meeting Abstract ORGN Super Taxa Animalia ORGN Organism Name C2-C12 cell line (Animalia): myoblast-osteoblast precursor cell line, osteoblast differentiation ORGN Organism Superterms Animals L193 ANSWER 5 OF 8 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 2000:413041 BIOSIS ΑN DN PREV200000413041 Specific inhibitors of the chymotryptic component of the ΤT proteasome are potent bone anabolic agents in vivo. Garrett, I. R. (1); Gutierrez, G. (1); Chen, D. (1); ΑU Rossini, G. (1); Escobedo, A. (1); Esparza, J. (1); Horn, D. (1); Crews, C. M.; Mundy, G. R. (1) (1) OsteoScreen, Inc., San Antonio, TX USA CS Journal of Bone and Mineral Research, (September, 2000) Vol. 15, No. SO Suppl. 1, pp. S197. print. Meeting Info.: Twenty-Second Annual Meeting of the American Society for Bone and Mineral Research Toronto, Ontario, Canada September 22-26, 2000 American Society for Bone and Mineral Research . ISSN: 0884-0431. DT Conference English LA English SLCytology and Cytochemistry - Animal \*02506 CC Biochemical Studies - General \*10060 Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and Biochemistry \*18004 General Biology - Symposia, Transactions and Proceedings of Conferences, Congresses, Review Annuals \*00520 BC Muridae 86375 ITMajor Concepts Biochemistry and Molecular Biophysics; Skeletal System (Movement and Parts, Structures, & Systems of Organisms TΤ bone: formation, skeletal system; osteoblast: proliferation, skeletal system Chemicals & Biochemicals IT potent bone anabolic agent: in-vivo; specific chymotryptic proteasome component inhibitor; statins Miscellaneous Descriptors IT Meeting Abstract ORGN Super Taxa Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name murine (Muridae) ORGN Organism Superterms Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates; Rodents; Vertebrates L193 ANSWER 6 OF 8 MEDITNE

1998312293

AN

MEDLINE

- gitomer 09 / 695807 98312293 PubMed ID: 9648487 DN TIHyperparathyroidism and its management. ΑU Sugimoto T Department of Medicine, Kobe University School of Medicine. CS NIPPON RINSHO. JAPANESE JOURNAL OF CLINICAL MEDICINE, (1998 Jun) SO 56 (6) 1591-7. Ref: 36 Journal code: KIM; 0420546. ISSN: 0047-1852. CY Japan Journal; Article; (JOURNAL ARTICLE) DT General Review; (REVIEW) (REVIEW LITERATURE) LA Japanese Priority Journals FS EM199809 ED Entered STN: 19980917 Last Updated on STN: 19980917 Entered Medline: 19980908 Hyperparathyroidism (HPT), resulting from the excess of endogenous ΑB parathyroid hormone is cited as one of diseases which cause secondary osteoporosis. HPT consists of primary (1 degree) and secondary (2 degrees) HPT, resulting mainly from chronic renal failure (CRF). HPT is easily distingishable from primary osteoporosis by biochemical measurements. Parathyroidectomy (PTX) is the only option available for the radical cure of 1 degree HPT and more than 10% increase in bone mass occurs after PTX. On the other hand, dietary phosphorus restriction, phosphorus binders, active vitamin D3 metabolites are useful for 2 degrees HPT due to CRF. When these treatments are not effective to inhibit PTH secretion adequately, oral active vitamin D3 pulse therapy, PTX and percutaneous ethanol injection therapy should be considered. CTCheck Tags: Human \*Hyperparathyroidism: CO, complications \*Hyperparathyroidism: TH, therapy Hyperparathyroidism, Secondary: CO, complications \*Osteoporosis: ET, etiology L193 ANSWER 7 OF 8 MEDLINE 96302997 MEDLINE AN DN 96302997 PubMed ID: 8741178 ΤI Suppressive effect of N-(benzyloxycarbonyl)-L-phenylalanyl-L-tyrosinal on bone resorption in vitro and in vivo. Woo J T; Yamaquchi K; Hayama T; Kobori T; Sigeizumi S; Sugimoto K; Kondo ΑU K; Tsuji T; Ohba Y; Tagami K; Sumitani K Sagami Chemical Research Center, Kanagawa, Japan. CS EUROPEAN JOURNAL OF PHARMACOLOGY, (1996 Apr 4) 300 (1-2) 131-5. SO Journal code: EN6; 1254354. ISSN: 0014-2999. CY Netherlands Journal; Article; (JOURNAL ARTICLE) DΤ LA English FS Priority Journals EM199610 Entered STN: 19961025 ED Last Updated on STN: 19961025 Entered Medline: 19961017 AB
- AB The suppressive effect of N-(benzyloxycarbonyl)-L-phenylalanyl-L-tyrosinal on bone resorption was examined in vitro and in vivo. This synthetic peptidyl aldehyde was found to be a potent and selective cathepsin L inhibitor in our screening for cysteine protease inhibitors. In the pit formation assay with unfractionated rat bone cells, 1.5 nM of this compound markedly inhibited parathyroid hormone-stimulated osteoclastic bone resorption. In addition, intraperitoneal administration of this peptidyl aldehyde (2.5-10 mg/kg) for 4 weeks suppressed bone weight loss dose dependently in the ovariectomized mouse,

experimental model of osteoporosis. Hydroxyproline measurement of the

decalcified femurs from these ovariectomized mice suggested that this compound acts as a bone resorption suppressor through the inhibition of collagen degradation. CT Check Tags: Animal; Female; Human \*Bone Resorption: PP, physiopathology \*Bone and Bones: DE, drug effects Bone and Bones: ME, metabolism \*Cathepsins: AI, antagonists & inhibitors \*Cysteine Proteinase Inhibitors: PD, pharmacology \*Dipeptides: PD, pharmacology Leucine: AA, analogs & derivatives Leucine: PD, pharmacology Mice Ovariectomy Rats Rats, Sprague-Dawley RN 66701-25-5 (E 64); 7005-03-0 (Leucine) CN O (Cysteine Proteinase Inhibitors); O (Dipeptides); O (N-(benzyloxycarbonyl)-phenylalanyl-tyrosinal); EC 3.4.- (Cathepsins); EC 3.4.22.15 (cathepsin L) L193 ANSWER 8 OF 8 MEDLINE 83293098 MEDLINE AN DN 83293098 PubMed ID: 6310016 Studies on osteoporoses. XI. Effects of a methylxanthine derivative. A TI preliminary report. ΑU Robin J C; Ambrus J L SO JOURNAL OF MEDICINE, (1983) 14 (2) 137-45. Journal code: IYG; 7505566. ISSN: 0025-7850. CY United States DT Journal; Article; (JOURNAL ARTICLE) English LA FS Priority Journals EM198310 ED Entered STN: 19900319 Last Updated on STN: 19900319 Entered Medline: 19831021 Heparin (500 U/kg s.c. B.I.D.) induced significant osteoporosis in AB C3H/St(Ha) female mice after 3 months of treatment. Pentoxifylline (12 mg/kg i.m. B.I.D.) prevented this experimental osteoporosis. Osteoporosis was measured by in vivo neutron activation analysis and results were confirmed by atomic absorption spectroscopy. Pentoxifylline (0.1-100 microgram/ml) increased calcium uptake and cAMP production in osteoblast-like bone cells isolated from fetal Sprague-Dawley rats. Theoretical implications for osteoblast control of bone resorption are discussed. CTCheck Tags: Animal; Female Bone Resorption Calcium: ME, metabolism Cyclic AMP: ME, metabolism Heparin Mice Mice, Inbred C3H Neutron Activation Analysis Osteoblasts: DE, drug effects Osteoblasts: ME, metabolism Osteoporosis: CI, chemically induced \*Osteoporosis: PC, prevention & control \*Pentoxifylline: TU, therapeutic use Rats Rats, Inbred Strains Spectrophotometry, Atomic Absorption

gitomer - 09 / 695807 Stimulation, Chemical \*Theobromine: AA, analogs & derivatives 60-92-4 (Cyclic AMP); 6493-05-6 (Pentoxifylline); 7440-70-2 RN (Calcium); 83-67-0 (Theobromine); 9005-49-6 (Heparin) => fil wpix FILE 'WPIX' ENTERED AT 16:31:59 ON 26 FEB 2002 COPYRIGHT (C) 2002 DERWENT INFORMATION LTD <20020221/UP> FILE LAST UPDATED: 21 FEB 2002 200212 MOST RECENT DERWENT UPDATE <200212/DW> DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE SDI'S MAY BE RUN ON EVERY UPDATE OR MONTHLY AS OF JUNE 2001. (EVERY UPDATE IS THE DEFAULT). FOR PRICING INFORMATION SEE HELP COST <<< >>> FOR UP-TO-DATE INFORMATION ABOUT THE DERWENT CHEMISTRY RESOURCE, PLEASE VISIT http://www.derwent.com/chemistryresource/index.html <<< >>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://www.derwent.com/dwpi/updates/dwpicov/index.html <<< => d all abeq tech tot L215 ANSWER 1 OF 2 WPIX COPYRIGHT 2002 DERWENT INFORMATION LTD 2000-686989 [67] WPIX DNC C2000-208928 Identifying a compound effective in treating multiple myeloma and myeloma TТ bone disease, involves subjecting the compound to an assay determining its ability to inhibit NF-kB or proteasomal activity. DC B04 IN MUNDY, G R (OSTE-N) OSTEOSCREEN INC CYC WO 2000061167 A2 20001019 (200067) \* EN PΙ 22p A61K038-04 RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE W: AU CA JP AU 2000042040 A 20001114 (200108) A61K038-04 A2 20020109 (200205) EN EP 1169049 A61K038-04

PA

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

WO 2000061167 A2 WO 2000-US9121 20000407; AU 2000042040 A AU 2000-42040 20000407; EP 1169049 A2 EP 2000-921764 20000407, WO 2000-US9121 20000407

AU 2000042040 A Based on WO 200061167; EP 1169049 A2 Based on WO 200061167 PRAI US 1999-289229 19990409

IC ICM A61K038-04

> A61K031-166; A61K031-40; A61P019-08 ICS

WO 200061167 A UPAB: 20001223 AΒ

> NOVELTY - Identifying a compound (I) effective in treating myeloma bone disease involves subjecting the compound to an assay to determine its ability to inhibit transcription factor NF-kB activity or production, or to an assay to determine its ability to inhibit proteasomal enzyme activity or production.

> DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a pharmaceutical composition for treating myeloma bone disease comprising (I); and
- (2) a method of treating myeloma bone disease by the administration of (I).

ACTIVITY - Osteopathic; cytostatic.

Nine C57BL/KaLwRij mice were inoculated with 0.5 asterisk 106 5TGM-1 cultured myeloma cells and tumor volume was assessed by the formula Tumor volume (cm3) = 4/3((length + width)-1)/2. The mice with tumors were randomized into two groups and treatment was commenced on day 35. One group has **PSI** injected directly into the tumors and the other group has only vehicle injected into the tumors. The tumors in the latter group (untreated mice) continued to grow, resulting in the mice dying between 42 and 55 days after myeloma cell inoculation. The size of the tumors in the treated mice decreased markedly and the mice remained healthy up to 3 months after tumor inoculation, even though treatment was discontinued. The result showed that the treated mice were alive and well with no signs of tumor 4 months after treatment.

MECHANISM OF ACTION - Inhibitor of NF-kB activity; inhibitor of proteasomal activity.

(I) reduces myeloma tumor volume, delays onset of limb paralysis, decreases the viability of myeloma cells and reduces the volume of tumor marker, IbG2b. (claimed).

USE - (I) is useful for treating multiple myeloma such as osteopenia, osteolytic lesions, osteopetrosis, bone fracture and osteolytic bone disease, and myeloma bone disease (claimed). Dwg.0/6

FS CPI

FA AB; DCN

MC CPI: B04-C01A; B10-A06; B10-A12A; B14-H01; B14-H01A; B14-L06

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AN 2000-171065 [15] WPIX

DNC C2000-053186

TI Compound that inhibits the activity of NF-kappa B useful for enhancing bone formation.

DC B04 B05

IN GARRETT, I R; MUNDY, G R; ROSSINI, G

PA (OSTE-N) OSTEOSCREEN; (OSTE-N) OSTEOSCREEN INC

CYC 73

PI WO 2000002548 A2 20000120 (200015)\* EN 37p A61K031-00

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW

W: AL AM AU BA BB BG BR CA CN CU CZ EE GE HU IL IN IS JP KP KR LC LK LR LT LV MD MG MK MN MX NO NZ PL RO SD SG SI SK TR TT US UZ VN

AU 9963109 A 20000201 (200028) A61K031-00

EP 1096924 A1 20010509 (200128) EN A61K031-00

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

ADT WO 2000002548 A2 WO 1999-US15533 19990709; AU 9963109 A AU 1999-63109

19990709; EP 1096924 A1 EP 1999-933827 19990709, WO 1999-US15533 19990709

FDT AU 9963109 A Based on WO 200002548; EP 1096924 Al Based on WO 200002548 PRAI US 1998-113947 19980710

IC ICM A61K031-00

AB WO 200002548 A UPAB: 20000323

NOVELTY - Enhancing bone formation, treating pathological dental conditions, treating degenerative joint conditions by administration of NF-kappa B inhibitor.

DETAILED DESCRIPTION - Enhancing bone formation or treating pathological dental conditions or treating degenerative joint conditions in a vertebrate animal comprises administration of a compound that inhibits the activity of NF-kB or that inhibits proteasomal activity or that inhibits production of proteasome proteins.

INDEPENDENT CLAIMS are included for the following:

- (1) treatment of a condition benefited by stimulating hair growth comprising administration of a compound that inhibits the activity of NF-kB or that inhibits **proteasomal** activity or that inhibits production of these proteins, and
- (2) identifying a compound which enhances bone growth or stimulates hair growth comprising subjecting a candidate compound to an assay to

assess its ability to inhibit:

- (a) NF-kB activity, or
- (b) the production of NF-kB, or
- (c) proteasomal activity, or
- (d) the production of enzymes with **proteasomal** activity, where for all the inhibitory compound is identified as a compound that enhances bone growth.

ACTIVITY - Osteopathic; Endocrine-Gen.; Screening; Vulnerary. PSI (N-carbobenzoyl-Ile-Glu-(OtBu)-Ala-Leu-CHO) was assayed in vitro for calvarial bone growth. Administered at 0.1, 1 and 5 mg/kg/day, the % increase in bone area compared to control was 21.7, 35.4 and 32.1%, respectively. The 1 and 5 mg/kg/day doses produced an increase in new bone width of 19.9%.

MECHANISM OF ACTION - Antimetastatic; Nuclear-Factor-Inhibitor-Kappa-B.

USE - The method can be used for enhancing bone formation, treating pathological dental conditions, degenerative bone conditions, osteoporosis, bone fracture or deficiency, primary or secondary hyperparathyroidism, periodontal disease or defect, metastatic bone disease, osteolytic bone disease, post-plastic surgery, post-prosthetic joint surgery, and post-dental implantation, and for stimulating hair growth (claimed). The compounds may also be useful in wound healing or tissue repair.

ADVANTAGE - None given.

Dwg.0/1

FS CPI

FA AB; DCN

MC CPI: B04-C01; B06-D13; B06-F05; B07-A02B; B07-D03; B10-A06; B10-A10;

B10-D02; B11-C08; B12-K04A; B14-D03; **B14-N01**;

B14-N06; B14-N11; B14-N17B; B14-R02

TECH

UPTX: 20000323

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Method: The compound does not inhibit the isoprenoid pathway. The compound is lactacystin, a peptidyl aldehyde or PTX. The method further comprises administration of one or more agents that promote bone growth or that inhibit bone resorption such as bone morphogenetic factors, anti-resorptive agents, osteogenic factors, cartilage-derived morphogenetic proteins, growth hormones, estrogens, bis phosphonates, statins or differentiating factors.